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obstructive pulmonary disease (COPD), a urological disorder such as neuropathy, incontinence or interstitial cystitis, or an inflammatory disorder.

According to another aspect of the invention there is provided a method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR, preferably hVR1 or hVR3, activity in a human patient which comprises administering to said patient an effective amount of a compound identified by the method referred to above. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), respiratory disorders such as asthma and chronic obstructive pulmonary disease (COPD) and urological disorders including diabetic neuropathy, incontinence and interstitial cystitis and inflammatory disorders.

According to another aspect of the invention there is provided a method of producing an hVR protein as hereinbefore described or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, comprising introducing into an appropriate cell line a suitable vector comprising a nucleotide sequence encoding for an hVR protein or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, under conditions suitable for obtaining expression of the hVR protein or a variant thereof, preferably hVR1 or hVR3 or a variant thereof.

Brief Description of the figures

Figure 1 is an alignment of hVR1 *in silico* derived clusters with rat VR1.

Figure 2 displays the human VR1 nucleotide sequence including the 5'UTR (nt – 773 to nt 0), coding region (nt 1 to 2517) and 3'UTR (nt 2518 to nt 3560) -- (SEQ ID NO:1)--.

Figure 3 illustrates the nucleotide and encoded amino acid sequence of the human VR1sequence -- (SEQ ID NO: 1 and SEQ ID NO:2) --.

Figure 4 depicts the amino acid sequence –(SEQ ID NO:2)-- of the hVR1 gene, the shading denotes predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed). The predicted phosphorylation sites are underlined. Figure 5 is a comparison of the amino acid sequences of the rat (rVR1) -- (SEQ

ID NO:3) -- and human (hVR1) vanilloid receptors -- (SEQ ID NO:2) --.

Figure 6 illustrates constructs pBluescriptSK(+) (A) and pCIN5-new (B) with the full length hVR1 gene cloned via Notl and EcoRI restriction sites.

Figure 7 shows a Slot Blot hybridisation with hVR1 probe with positive labelling of both rat and human DRG mRNA.

Figure 8 displays a Western blot probed with anti-VR1 antibodies with the arrow indicating the VR1 specific protein.

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Figure 9 shows localisation of VR1 in rat DRG tissue sections, the arrow points to VR1 expressing small diameter (<25µn) neurone cell bodies.

Figure 10 depicts the *in situ* localisation of VR1 in human DRG sections (A) and human skin (B).

Figure 11 illustrates the functional response to capsaicin and blockade by capsazepine (CPZ) (A) with the current voltage relationship plotted in (B) on human VR-1 channels, transiently expressed in HEK293T cells.

Figure 12 shows capsaicin-induced desensitisation of human VR-1 channels in the presence of 2mM external calcium (A), maximum current (65mV) against time (B) and current voltage relationship in the absense of Ca²⁺ (C).

Figure 13 shows the influx of calcium into transiently transfected HEK293T cells over a time course in the presence of agonist capsaicin, anandamide and resiniferatoxin in the absence (A, B, D and F) or presence (C, E, G) of the antagonist, capsezipine.

Figure 14 illustrates a graphical presentation the results shown in figure 13 examining the response of hVR1 transfected HEK293T cells over time before and after exposure to agonists: capsaicin, anandamide and resiniferatoxin in the absence (A, B, D and F) or presence (C, E, G) of the antagonist, capsezipine.

Figure 15 displays the proposed assay strategy to carry out drug screening. Figure 16 displays an alignment of *in silico* derived hVR3 specific clusters with rat VR1.

Figure 17 depicts the hVR3 nucleotide sequence including the 5' UTR (nt –686 to nt 0) Coding region (nt1 to nt 2889), 3'UTR (nt 2890 to nt 3418) -- (SEQ ID NO:4) --.

Figure 18 shows the nucleotide and amino acid sequence of hVR3 -- (SEQ ID NO:4 and SEQ ID NO:5) --.

Figure 19 is of the amino acid sequence of hVR3 -- (SEQ ID NO:5) --, the shading denotes predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed).

Figure 21 illustrates a multiple comparison of the amino acid sequences of the rat VR1 --(SEQ ID NO:3) -- and the human vanilloid receptors: hVR1, hVRL-1 and hVR3 -- (SEQ ID NO:2), (SEQ ID NO:6) and (SEQ ID NO:5) --, respectively. Figure 22 Northern Blot hybridisation with hVR3 probe with strong signals detected in trachea (A), prostate (B), placenta, kidney and pancreas (C).

Detailed Description of the Invention

Throughout the present specification and the accompanying claims the words "comprise" and "include" and variations such as "comprises", "comprising", "includes" and "including" are to be interpreted inclusively. That is, these words are intended to convey the possible inclusion of other elements or integers not specifically recited, where the context allows.

As referred to above, the present invention relates to isolated human vanilloid receptor (hVR) proteins, and in particular to the human vanilloid receptors which will be termed respectively human vanilloid receptors 1 and 3 (hVR1, and hVR3), sequence information for which is provided in figures 2 (hVR1) and 17 (hVR3). In the context of this invention the term "isolated" is intended to convey that the receptor protein is not in its native state, insofar as it has been purified at least to some extent or has been synthetically produced, for example by recombinant methods. The term "isolated" therefore includes the possibility of the receptor protein being in combination with other biological or non-biological material, such as cells, suspensions of cells or cell fragments, proteins, peptides, organic or inorganic solvents, or other materials where appropriate, but excludes the situation where the receptor protein is in a state as found in nature.

Routine methods, as further explained in the subsequent experimental section, can be employed to purify and/or synthesise the receptor proteins according to the invention. Such methods are well understood by persons skilled in the art, and include techniques such as those disclosed in Sambrook, J. et al. (28), the disclosure of which is included herein in its entirety by way of reference.

By the term "variant" what is meant throughout the specification and claims is that other peptides or proteins which retain the same essential character of the human vanilloid receptor proteins for which sequence information is provided, are also intended to be included within the scope of the invention. For example,

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The compounds may be administered via enteral or parenteral routes such as via oral, buccal, anal, pulmonary, intravenous, intraarterial, intramuscular, intraperitoneal, topical or other appropriate administration routes.

5 The present invention will be further explained, by way of examples, in the appended experimental section. Reference examples are provided.

Experimental details

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10 Reference Example A: Identification of related human ESTs (Expressed Sequence Tags) (19) to the rat VR1 sequence by in silico analysis

The full-length rat VR1 amino acid sequence (15) was used as a query sequence using the tBlastn (20) alignment program to identify related human genes in the dbEST (21) and Incyte (Incyte Pharmaceuticals, Inc., 3174 Porter Drive, Palo Alto, California 94304, USA) databases. Several human ESTs were identified and those with similarities greater than 50% selected for further analysis. One of these ESTs was T12251 previously shown to have 68% aminoacid identity and 84% similarity over a region of 70 amino acids (15). Full-length cloning and functional characterisation of the gene represented by this cluster has been completed (30). This gene was denoted hVRL-1 and encoded a protein of 764 amino acid protein -- (SEQ ID NO:6) -- that was 48 % identical to the rat VR1 protein. All human ESTs from both databases were clustered to identify overlapping identical ESTs belonging to the same transcript. The GCG package (Wisconsin Package Version 9.0, Genetics Computer Group (GCG), Madison, Wisconsin) and a program developed in house termed ESTBlast (22) were used to build up these clusters. In total, forty-three ESTs derived from different tissue sources and both EST databases were clustered into ten groups, one of these clusters represented hVRL-1. The remaining nine clusters have been named hVRa, hVRb, hVRc, hVRd, hVRe, hVRf, hVRg, hVRh and hVRi. For each EST the tissue source was assigned according to the annotations in the dbEST and Incyte databases. Since no obvious starting codon was present and the cluster sequences were shorter than the rat VR1 transcript none of these clusters were likely to represent a full-length vanilloid receptor transcript. Finally hVRq, hVRh and hVRi collapsed into a single contig. Sequence analysis

35 has shown that

Reference Example B2: Sequencing of cl nes

All DNA sequences were determined by automated DNA sequencing based on the dideoxy chain-termination method using the ABI 373A / 377 sequencers (Applied Biosystems). Sequence-specific primers were used with the 'Big-Dye' Terminator Cycle Sequencing kit (Applied Biosystems). The nucleotide sequence was analysed using programs from the University of Wisconsin Genetics Computer Group package.

More specifically when sequencing an EST clone, the following protocol was adopted. The EST1 clone was grown using standard procedures and DNA was isolated using Qiagen columns. SP6 (5' ATTTAGGTGACACTATAG) -- (SEQ ID NO:7) -- and T7 (5' TAATACGACTCACTATAGGG) -- (SEQ ID NO:8) -- primers flanking the cloning site were used to sequence both ends. Plasmid DNA (0.6 pmol) was used with 10.0 pmol of each primer for the dye terminator reaction. The SP6 end corresponded to the *in silico* derived EST sequence (identical to EST1). The T7 end did not have homologies with VR1 nor did it possess a long open reading frame or a polyadenylation motif. The size of the insert was determined by enzyme digestion of the DNA with the endonucleases Notl and EcoRI and calculated to be approximately 3kb.

Plasmid DNA (50ng) was used to amplify the insert by Polymerase Chain Reaction (PCR) with T7 and SP6 as primers. The PCR conditions included an initial hot-start at 94°C for 2 minutes, followed by 35 cycles at 94°C for 45 seconds, 50°C for 45 seconds and 72°C for 1 minute and terminated by 5 minutes at 72°C. The resulting PCR amplicon was separated on a 1.2% agarose gel and shown to be of ~3kb in size.

To fully sequence the PCR product the nuclease-Bal-31 technique was used where both strands of duplex DNA are degraded from both ends (23). After ethanol precipitation of the PCR product, the pellet was re-suspended in 30ml of 1X Bal-31 buffer (add buffer composition). A time-course digest with 2 units of Bal-31 enzyme (Roche Molecular Biochemicals) was carried out with 12 time points taken over 90 minutes (30 seconds, 1, 2, 3, 5, 7, 10, 15, 25, 45, 75 and 90 minutes). Three pools were made respectively from digests 1 to 4, 5 to 8 and 9

to 12. Each pool was blunt-ended and sub-cloned into the pCR-Script SK (+) plasmid from Stratagene at the Srfl site. After transformation, 16 colonies from each pool were screened by PCR with the flanking Reverse GGAAACAGCTATGACCATG) -- (SEQ ID NO:9) -- and M13-20 GTAAAACGACGCCAGT) -- (SEQ ID NO:10) -- primers. The amplicons of 6 positive colonies per pool were subjected to direct sequencing (24) using the T3 (5' AATTAACCCTCACTAAAGGG) -- (SEQ ID NO:11) -- and T7 primers. The DNA sequences obtained were assembled using the GCG package, translated and aligned against the rat VR1 gene using the Blast tools. After analysis, the 3079bp amplicon was shown to have 2 introns of 603bp and 1221bp. The latter intron was located at the 3'end of the PCR product. The coding sequence covered 1255 bp and was separated by the former intron. Therefore the clone EST1 was likely to be a partially spliced and incomplete cDNA.

The clone belonging to cluster 1b (EST3) and derived from a kidney cDNA library was ordered and sequenced using the Bal-31 technique. After assembly of the sequences using the GCG package an identical overlap was identified with the DNA sequence of the cluster hVRc. Moreover a 3'end with a polyadenlyation signal and tail was identified. The complete sequence of the combined hVRb Bal-31 derived sequence and hVRc was 2063 bp (1020 bp of coding and 1043 bp of 3' untranslated sequence).

Reference Example B3: Amplification of the middle section of hVR1 using the Polymerase Chain Reaction

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We formulated the hypothesis that both sequences (hVRa and hVRb/c) were part of a common transcript. If the human and rat VR1 were going to be similar, the 2 contigs should be separated by a gap of approximately 275bp. Primers were designed on both sides of the gap to amplify mRNA from brain tissues in order to clone the gap. A smear was obtained with the sense primer (5' TCTACTTCGGTGAACTGCCC) -- (SEQ ID NO:12) -- and antisense (5' ACGGCAGGGAGTCATTCTTC) -- (SEQ ID NO:13) --. For specificity 50ng of the PCR (5' product were amplified with the nested sense CTGCAGAACTCCTGGCAGA) -- (SEQ ID NO:14) -- and antisense (5' GTCACCACCGCTGTGGAAAA) -- (SEQ ID NO:15) -- primers. The 900bp nested amplicon was sequenced and shown to be identical to hVRa at one end 5

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hVRb/c at the other end. The middle part of the PCR product was homologous to the rat VR1 sequence. This region corresponded to 91 amino acids. When the sequences of hVRa, hVRb/hVRc and the internal amplicon are combined the total length of the Open Reading Frame (ORF) is 824 amino acids followed by a 3' untranslated sequence of 1043 bp. The human amino acid sequence is 87% identical to the rat sequence over that part of the coding region. This sequence was termed hVR1 because of its high degree of identity with the rat VR1 sequence.

10 Reference Example B4: Isolation of the 5' Terminus of hVR1 by PAC isolation

Since no start codon was identified at the 5' end an additional strategy was designed to identify the full-length sequence. Two primers, sense (5' TCCTCTGGCTTCCAACCCGTT) -- (SEQ ID NO:16) -- and antisense (5' GAACTGGGCAGAAAGTGCCT) -- (SEQ ID NO:17) -- were designed to amplify a 150bp product from the first intron mentioned in reference example B2. A P1 Artificial Chromosome (PAC) genomic clone (25) was isolated by PCR screening of a PAC library (Genome Systems, St Louis, Missouri). PAC DNA was recovered by using standard plasmid isolation protocol (26). An anti-sense primer was designed (5' CTGGAGTTAGGGTCTCCATCC) -- (SEQ ID NO:18) -to sequence the genomic clone towards the potential 5' end of the gene. An open reading frame with a starting codon was identified. The gene structure was confirmed by using the GenScan software (27). The complete gene has a nucleotide sequence of 2517bp (figure 2) and encoded a 839 amino acid protein (Figures 3 and 4). The gene was named hVR1. Multiple alignment of the amino acid sequence of hVR1 and rat VR1 shows a remarkable degree of identity and similarities between both sequences (figure 5). The rVR1 and hVR1 amino acid sequences are 86% identical. Moreover after protein analysis 6 trans-membrane domains and 3 ankyrin binding domains were identified in hVR1 as in the rat VR1 gene.

Example 1: Full-length Amplification of hVR1 from human DRG and assembly into cloning vectors

35 HVR1 was PCR amplified in three sections from human DRG template. The 5' fragment was amplified using a sense primer encoding a Notl site and a strong

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Kozak motif followed by specific gene sequence (5' GTCATAGCGGCCGCCGCCACCATGAAGAAATGGAGCAGCAC) -- (SEQ ID NO:19) -- and an antisense primer (5' AGGCCCACTCGGTGAACTTC) --(SEQ ID NO:20)--. The thermo-cycling conditions used for this amplification included a hot start at 94°C for 4 mins, followed by 35 cycles of 94°C for 1 min, 54°C for 1 min. and 72°C for 1 min. A final extension step of 72°C for 5 min completed the reaction. The resulting PCR products were separated on a 2% agarose gel and cloned into pCR®II-TOPO according to the manufacturers instructions supplied with the TOPO™ TA Cloning® kit (Invitrogen). The middle was PCR amplified using the sense primer: section of hVR1 GACGAGCATGTACAATGAGA -- (SEQ ID NO:21) -- and antisense primer: 5' GTCACCACCGCTGTGGAAAA -- (SEQ ID NO:22) --. The cycling conditions included a hot start at 94°C for 4 mins, followed by 35 cycles of 1 min at 94°C, 56°C and 72°C. A final extension step of 72°C for 5 min completed the reaction. A band of approximately 870 bp was excised from a 2 % agarose gel and cloned as detailed by the TOPO™ TA Cloning® kit into the vector pCR2.1®-TOPO. Finally the 3' end was PCR amplified with the sense primer: 5' TGTGGACAGCTACAGTGAGA -- (SEQ ID NO:23) -- and the antisense primer: 5'TGCACTGAATTCGAGCACTGGTGTTCCCTCAG -- (SEQ ID NO:24) -- which encoded an EcoRI site for cloning. The PCR conditions included a 90 sec hot start at 94°C followed by 35 cycles of 94°C for 50 sec, 50°C for 50 sec and 72°C for 50 sec. The cycling was completed with a 72°C step for 5 min. PCR products were separated on a 2% agarose gel and cloned into the vector pCR2.1®-TOPO.

Resulting clones for each of the three hVR1-fragments were taken for sequence analysis and separate clones coding a consensus sequence were used in the full length assembly of the gene. The Notl/Dralll (New England Biolabs) digested 5' end fragment ligated together with the middle Dralll/EcoRl fragment into a Notl/EcoRl restricted pBluescript SK (+) vector (Stratagene). Finally, the remaining 3' fragment was introduced into the resulting construct via Mscl and EcoRl restriction sites, a map of the resulting construct is displayed in figure 6A.

Several clones were selected for sequence analysis to confirm that constructs still encoded the hVR1 consensus sequence. These were then digested with Notl/EcoRl and ligated into the mammalian expression vector pClN5-new (a modified version of pClN1 (32) having an IVS deletion as well as a 36 bp

deletion repositioning the start codon of neomycin phosphotransferase immediately after the upstream EMVC IRES) as illustrated in figure 6B.

Example 2: Chromosomal Localisation

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The primers used to isolate the PAC clone (reference example B4) were selected for PCR on the G3 radiation hybrid panel from Stanford commercially available from Research Genetics (Huntsville, Alabama). The positive lanes and negative patterns were analysed using the public web server at Stanford University (http://www-sghc.stanford.edu). After analysis the hVR1 gene appears to be located on human chromosome 17 around marker SHGC-36073 (lod score=9.55).

Example 3: mRNA Distribution

The tissue distribution of hVR1 was established by slot-blot hybridisation. RNA was transferred onto a sheet of GeneScreen hybridisation transfer membrane (DUPONT) sandwiched in a slot blotter by suction via a vacuum pump. Once the membrane was rinsed in 2x SSC (3M sodium chloride and 0.3M sodium citrate pH7) for 2 min it was exposed to UV using an Ultraviolet crosslinker (Amersham Life Science) for 1min at 15000uW/cm² thus enabling cross-linkage of the RNA onto the membrane. The amounts of RNA on the blot are unknown. The probe was obtained by PCR amplification of a 260 bp product of the coding region of hVR1 with the following two primers: 5' TGTGGACAGCTACAGTGAGA -- (SEQ ID NO:25) -- and 5' GTGGAAAACCCGAACAAGA --(SEQ ID NO:26) --. Membranes were hybridised for 4 hr shaking at 60°C in a 10% dextran sulphate, 1% SDS (sodium dodecyl sulphate) and 1M NaCl solution. The probe was labelled with [α32P]dCTP (Amersham) using the Rediprime™DNA labelling system (Amersham), so as to obtain approximately 500,000cpm of the labelled probe per ml of prehybridisation solution. Briefly 100ng of probe was boiled for 3 minutes (denaturization) and then cooled on ice for 2 minutes in a total volume of 45µl. This was added to the labelling tube from the kit together with 3µl of 32P dCTP followed by an incubation at 37°C for 30 minutes. 400µl of Herring Sperm DNA (Sigma) at a concentration of 8µg/ml was added to the labelled probe and heated at 99°C for 3 minutes followed by rapid cooling on ice. The labelled probe was added and mixed well in pre-hybridisation solution. The membranes were hybridised overnight at 55°C.

The membranes were then washed, first at room temperature in 2xSSC and 1% SDS for 5 minutes, followed by 2x SSC and 1% SDS for 30 min at 50°C. If necessary further washes with 1x SSC and 0.5% SDS or 0.1xSSC and 0.1% for 30 mins at the same temperature were carried out. The membranes were then exposed to Scientific Imaging Film AR (Kodak) using intensifying screens at – 70°C overnight and the film developed.

The results are shown on figure 7. Strong signals were observed with the positive controls (slots 4B and 5B). Signals are detected on the human DRG slots (1A and 1B). No signals were detected with the water control (slot 3B). Three multi-tissue northern blots (Clontech) with a wide range of tissues have also been hybridised with the same probe, however no signals were detected. RT-PCR was performed on various tissues with the primer combination used to amplify the probe. A strong band was detected in DRG RNA. Taken together these hybridisations suggest that hVR1 is specifically expressed in neuronal tissue and DRG in particular.

Example 4: Design and production of Anti-hVR1 Antibody

The peptides CHIFTTRSRTRLFGKGDSEEASC -- (SEQ ID NO:27)-- (peptide68) and CGSLKPEDAEVFKDSMVPGEK -- (SEQ ID NO: 28)-- (peptide69) were synthesised by standard solid phase techniques and purified by gel filtration chromatography. These peptides were conjugated via their Cys residues to the carrier protein, Tuberculin PPD (purified protein derivative) using sulpho-SMCC (sulfosuccinimidyl 4-[N-maleimidomethyl]-cyclohexan-1-carboxylate). Rabbits, previously sensitised to Bacillus Calmette Guerin (BCG), were inoculated with the resulting conjugates emulsified in incomplete Freund's adjuvant at approx monthly intervals. Serum was prepared from blood samples taken 7 days after each immunisation. The specific antibody response was followed by indirect enzyme-linked immunosorbent assay (ELISA) using free peptide as antigen. Immunoglobulins were purified from high titre sera using immobilsed peptide affinity columns (sulpholink Pierce). Rabbits designated M143, 144 and 145 received peptide68 conjugate, rabbits M146, 147 and 148, peptide69 conjugate.

The antibodies have been validated by specific staining of the recombinant protein expressed in HEK293 cells. Whole cell lysates were prepared in Sample

peak heights are reduced in cells pre-incubated in CPZ. The same FLIPR assay may be used to monitor the response of human VR1 on exposure to agonists and antagonists.

5 Example 8: Example of a screen using human VR1.

FLIPR assay technology may be utilised to screen for hVR1 modulators according to the procedure described in figure 15. Human VR1 may be gated with protons, capsaicin or heat.

Reference Example C: Identification and partial characterisation of additional human vanilloid receptors (referenence examples C1-C3):

Reference Example C1: Identification and characterisation of a novel vanilloid–like receptor, hVR3

ESTs belonging to the remaining clusters were characterised by *in silico* cloning (reference example A). The following clones were used during this process: - EST6/EST7 (hVRd), -EST8. (hVRe), - EST9/EST10. (hVRf). These EST clusters have been aligned with rat VR1 in figure 16, note that this diagram is not to scale.

Reference Example C2: Sequencing of clones

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Further sequencing, as detailed in reference example B2, and *in silico* cloning, enabled clusters hVRd, hVRe and hVRf to collapse forming a single contig of 583 amino acids. This sequence was named hVR3 and has 49 % identity with the rat VR1 sequence. It was unlikely that this single contig was a full-length vanilloid receptor transcript as no obvious starting codon was present and it was shorter than the rat VR1 transcript.

Reference Example C3: Identification of the 5' terminus of hVR3

Two primers (sense primer 5' ATGGCCACCAGCAGGGTTAC -- (SEQ ID NO:29) -- and antisense primer 5' TCTGCCAGGTTCCAGCTG) -- (SEQ ID NO:30) -- designed to PCR amplify an amplicon stretching the 3' end of hVR3 and its 3'utr were used to isolate a genomic PAC clone (Genome Systems. St Louis, Missouri). The hVR3 specific PAC clone was then used as template to generate a library. This was achieved by sonicating 6μg of Qiagen purified PAC construct, size selecting fragmented DNA of 500-

2000bp. These resulting fragments were then blunt ended and cloned into the vector pCR®-Blunt as detailed in the manufacturers protocol supplied with the Zero Blunt™ PCR cloning kit (Invitrogen). Clones were then sequenced (reference example B2) to identify the complete 5' end of the hVR3 transcript. The full-length nucleotide sequence of the hVR3 gene is displayed in figure 17. Figure 18 illustrates both nucleotide and encoded amino acid sequence of the human VR1 and figure 19 depicts the amino acid sequence of the hVR3 gene with shaded regions denoting predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed).

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Example 9: Full-length Amplification of hVR3 from human kidney template

Human kidney was used as a source of template for the PCR amplification of hVR3. Primers used for amplification were designed to isolate the gene in three fragments. Primers designed to isolate the 5' end included a sense primer encoding a Notl site and a strong Kozak motif followed by gene specific sequence (5' GTCATAGCGGCCGCGCGCCACCATGCCCAGGGTAGTTGGAC -- (SEQ ID NO: 31) -- and antisense primer (5' CACCTCTTGTTGTCACTGGA) --(SEQ ID NO:32) --. The PCR conditions used were a hot start at 94°C for 4 mins, followed by 35 cycles of 94°C for 1 min, 56°C for 1 min and 72°C for 1 min and finally one cycle at 72°C for 5 min. The resulting PCR products were separated on a 2% agarose gel and cloned into pCR®II-TOPO according to the manufacturers instructions supplied with the TOPOTM TA Cloning® kit (Invitrogen). The middle fragment was PCR generated using sense and antisense primers 5' CAAATCTGCGCATGAAGTTCCAG -- (SEQ ID NO:33) -and 5' GCCACGAGAAGTTCCACGTAGTG -- (SEQ ID NO:34) -- respectively in the presence of 5% DMSO. PCR thermo-cycling required 35 cycles of 1 min at 94°C, 58°C and 72°C for successful amplification of the fragment which was then excised from a 2% agarose gel for cloning into the pCRII®-TOPO vector. Finally the 3' amplified with sense primer 5' fragment was GCTGCTCCCATTCTTGCTGA -- (SEQ ID NO:35) -- and an antisense primer 5' TGCACTCTCGAGAAATGAGTGGGCAGAGAAGC -- (SEQ ID NO:36) -encoding a Xhol restriction site. This fragment was successfully amplified using a hot start at 94°C for 4 min followed by 35 cycles of 94°C for 50 sec, 48°C for 50 sec and 72°C for 2 min. The cycling was completed with a 72°C step for 5 min. The amplified fragment was excised from a 2% agarose gel and clone into the pCRII®-TOPO vector.

Resulting clones for each of the three PCR generated hVR3-fragments were taken for sequence analysis and separate clones coding a consensus sequence were used in the full-length assembly of the gene. The DrallI restriction site of the pBluescript SK (+) vector (Stratagene) was firstly abolished by digestion with DralII followed by a blunt ending step using T₄ DNA polymerase (New England Biolabs). This modified vector was then restricted to enable the ligation of both a Notl/Ncol 5' fragment and Ncol/ EcoRI middle fragment. Finally, the remaining 3' fragment was introduced into the resulting construct via DralII and Xhol sites (figure 20A).

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Several clones were selected for sequence analysis to confirm that the constructs still encoded the hVR3 consensus sequence. These were then digested with Notl/XhoI and ligated into the mammalian expression vector pCDNA3.1 (+) (Invitrogen) as seen in figure 20B. The resulting hVR3 consensus sequence is shown in the multiple alignment along with the full-length sequence of hVR1 and the published hVRL-1 in figure 21.

Example 10: Chromosomal localisation

The 3' terminus, including the 3' UTR sequence of hVR3 was used to design two primers amplify product of 360 bp: sense primer ATGGCCACCAGCAGGGTTAC -- (SEQ ID NO:37) -- and antisense primer 5' TCTGCCAGGTTCCAGCTG -- (SEQ ID NO:38) --. The G3 radiation hybrid panel from Stanford University (Research Genetics, Huntsville, Alabama) was screened by PCR. The positive and negative lanes were analysed using the public web server at Stanford University (http://www-sghc.stanford.edu). After analysis the hVR3 gene appears to be located on human chromosome 12 around markers D12S177E (lod score=15) and D12S1893 (lod score=14).

Example 11: mRNA distribution

The following primers (5' ACAAGAAGGCGGACATGCGG -- (SEQ ID NO:39) -- and 5' ATCTCGTGGCGGTTCTCAAT) -- (SEQ ID NO:40) -- were used to obtain a PCR product from the coding region of hVR3. This amplicon was used as a probe on multi-tissue northern blots, the protocol of which is detailed in example 3, to determine the tissue distribution of the gene (figures 22A, 22B and 22C). A transcript of approximately 3.8 kb was detected in the following tissues (the intensities of the

FIG. 2

hVR1 SEQUENCE INCLUDING THE 5'UTR (nt -773 TO nt 0), CODING REGION (nt 1 TO 2517) AND 3'UTR (nt 2518 TO nt 3560) -- (SEQ ID NO:1) --

-773		-714
-713	aaggccagaagcttgacagatgttgattcataaaaatgcaaaagccaaaatccaaaatct	-654
-653		-594
-593		-534
-533	ccagtetetgccgctcaccctattccagggacacagtetgcttggctcttctggactgag	-474
-473	ccatcctcatcaccgagatcctccctgaattcagcccacgacagccaccccggccgtttt	-414
-413	ccttgttctgtgtgggaagggaggcagcgggtggttatcaacctcaccctgcagaggag	-354
-353	gcacctgaggcccagagacgagggggatgggtctaacccagaaccacagatggctctga	-294
-293	geegggggeetgteeaceeteecaggeegaegteagtggeegeaggaetgeetgggeeet	-234
-233	getaggeetgeteacetetgaggeetetggggtgagaggttcagtcetggaaacaettca	-174
-173	gttctagggggctgggggcagcagcaagttggagttttggggtaccctgcttcacagggc	-114
-113	cettggcaaggaggcaggtggggtctaaggacaagcagtccttactttgggagtcaacc	-54
-53	ccggcgtggtggctgctgcaggttgcacactgggccacagaggatccagcaaggATGAAG	6
7	AAATGGAGCAGCACAGACTTGGGGGCAGCTGCGGACCCACTCCAAAAGGACACCTGCCCA	66
67	GACCCCTGGATGGAGACCCTAACTCCAGGCCACCTCCAGCCAAGCCCCAGCTCTCCACG	126
127	GCCAAGAGCCGCACCCGGCTCTTTGGGAAGGGTGACTCGGAGGAGGCTTTCCCGGTGGAT	186
187	TGCCCTCACGAGGAAGGTGAGCTGGACTCCTGCCCGACCATCACAGTCAGCCCTGTTATC	246
247	ACCATCCAGAGGCCAGGAGACGGCCCCACCGGTGCCAGGCTGCTGTCCCAGGACTCTGTC	306

FIG. 3

NUCLEOTIDE AND AMINO ACID SEQUENCE OF hVR1 INCLUDING 44 (SEQ ID NO:1) -- THE 5'UTR (nt -773 TO nt 0), CODING REGION (nt TO 2517) AND 3'UTR (nt 2518 TO nt 3560) -- (SEQ ID NO:2) --

	,	
-773	ccccagccacacacacacacacacacacacacacacaca	-714
-713	aaggccagaagcttgacagatgttgattcataaaaatgcaaaagccaaaatccaaaatct	-654
-653	tgtataagctcagtggctgtggcagcgaggttgaagagcaaaggcaggc	-594
-593	ctgatgatgtgtggacccgttgcacagcagggcccgcagtgcggtgtggggtgtgggg	-534
-533	ccagtctctgccgctcaccctattccagggacacagtctgcttggctcttctggactgag	-474
-473	ccatcctcatcaccgagatcctccctgaattcagcccacgacagccaccccggccgtttt	-414
-413	ccttgttctgtgtgggaagggaggcagcgggtggttatcaacctcaccctgcagaggag	-354
-353	gcacctgaggcccagagaggagggatgggtctaacccagaaccacagatggctctga	-294
-293	gccgggggcctgtccaccctcccaggccgacgtcagtggccgcaggactgcctgggccct	-234
-233	gctaggcctgctcacctctgaggcctctggggtgagaggttcagtcctggaaacacttca	-174
-173	gttctagggggctgggggcagcagcaagttggagttttgggggtaccctgcttcacagggc	-114
-113	ccttggcaaggagggcaggtggggtctaaggacaagcagtccttactttgggagtcaacc	-54
-53 1	ccggcgtggtggctgctgcaggttgcacactgggccacagaggatccagcaaggATGAAG M K	6 2
7	AAATGGAGCAGCACAGACTTGGGGGCAGCTGCGGACCCACTCCAAAAGGACACCTGCCCA	66
3	K W S S T D L G A A A D P L Q K D T C P	22
67	GACCCCTGGATGGAGACCCTAACTCCAGGCCACCTCCAGCCAAGCCCCAGCTCTCCACG	126
23	D P L D G D P N S R P P P A K P Q L S T	42
127	GCCAAGAGCCGCACCCGGCTCTTTGGGAAGGGTGACTCGGAGGAGGCTTTCCCGGTGGAT	186
43	A K S R T R L F G K G D S E E A F P V D	62
187	TGCCCTCACGAGGAAGGTGAGCTGGACTCCTGCCCGACCATCACAGTCAGCCCTGTTATC	246
63	C P H E E G E L D S C P T I T V S P V I	82
247	ACCATCCAGAGGCCAGGAGACGCCCCACCGGTGCCAGGCTGCTGTCCCAGGACTCTGTC	306
83	T I Q R P G D G P T G A R L L S Q D S V	102
307	GCCGCCAGCACCGAGAAGACCCTCAGGCTCTATGATCGCAGGAGTATCTTTGAAGCCGTT	366
103	A A S T E K T L R L Y D R R S I F E A V	122
367	GCTCAGAATAACTGCCAGGATCTGGAGAGCCTGCTGCTCTTCCTGCAGAAGAGCAAGAAG	
123		426
		142
427	CACCTCACAGACAACGAGTTCAAAGACCCTGAGACAGGGGAAGACCTGTCTGCTGAAAGCC	486
143	HLTDNEFKDPETGKTCLLKA	162
487	ATGCTCAACCTGCACGACGGACAGAACACCACCATCCTCCTCCTCCTCCTCCACATCCCCCCC	516

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FIG. 4

		Aiviti	IO ACID SE	QUENCE O	1141/1(2	EQ ID NO:2).	
	1	MKKWSSTDLG	AAADPLQKDT	CPDPLDGDPN	SRPPPAKPQL	STAKSRTRLF	
	51	GKGDSEEAFP	VDCPHEEGEL	DSCPTITVSP	VITIQRPGDG	PTGARLLSQD	
	101	SVAASTEKTL	RLYDRRSIFE	AVAQNNCQDL	ESLLLFLQKS	kkhl i dnefk	
	151	DPETGKTCLL	KAMLNLHDGQ	NTTIPLLLEI	ARQTDSLKEL	VNASYTDSYY	
	201	KGQTALHIAI	ERRNMALVTL	LVENGADVQA	AAHGDFFKKT	KGRPGFYFGE.	
	251	LPLSLAACTN	QLGIVKFLLQ	NSWQTADISA	RDSVGNTVLH	ALVEVADNTA	
	301	DNTKFVTSMY	NEILILGAKL	HPTLKLEELT	NHKGMTPLAL	AAGTGKIGVL	
	351	AYILQREIQE	PECRHLSRKF	T EWAYGPVHS	SLYDLSCIDT	CEKNSVLEVI	
	401	AYSSSETPNR	HDMĹLVEPLN	RLLQDKWDRF	VKRIEVENEL	VYCLYMIIFT	
	451	MAAYYRPVDG	LPPFKMEKIG	DYFRVTGEI	SVLGGVYFFF	RGIOXFLORR	
	501	PSMKTLFVI S	YSEMEFFLQS;	LFMLATVVLY	es ilkeyvas	MVFSLALGWT	
	551	NMLYYTRGFQ	OMGIYAVMI	KMILRULCRE	MEVYIVELEG	FSTAVVTLIE	
	601	DGKNDSLPSE	STSHRWRGPA	CRPPDSSYNS	LYSTCLELFK	FTIGMGDLEF	
	651	TENYDEKAVE:	TILLLAYVIL	TYTELLNMGI:	ALMGETVNKI	AQESKNI W KL	
	701	QRAITILDTE	KSFLKCMRKA	FRSGKLLQVG	YTPDGKDDYR	WCFRVDEVNW	
	751	TTWNTNVGII	NEDPGNCXGV	KRTLSFSLRS	SRVSGRHWKN	FALVPLLREA	
	801	SARDRQSAQP	EEVYLRQFSG	SLKPEDAEVF	KSPAASGEK*		
Key							
T/S predicted phosphorylation sites							
Transmembrane domains							
	Ankyrin binding domains						

11/41 FIG. 5

COMPARISON OF THE AMINO ACID SEQUENCE OF THE RAT (VR1)--(SEQ ID NO:3)-- AND HUMAN (hVR1) VANILLOID PROTEINS.--(SEQ ID NO:2)--

		(_ ,
	10	20	30	40	50
VR1	MEQRASLDSEESE	And the second s		Grant Francisco I. Proces	
hVR1	MKKWSSTDLGAAA	DPLQKDTCPDPI	DGDPNSRPPI	PARPOLSTAK	SRTRLF
	60	70	80	90	100
VR1	GKGDSEEASPLDC				
hVR1	GKGDSEEAFPVDC	PHEEGELDSCPT	TITVSPVITI	PREGOGPTGA	RLLSQD
	110	120	130	140	150
VR1	SVSAG.EKPPRLY				
hVR1	SVAASTEKTLRLY	Drrsifeavaqn	INCODLESLLI	LFLQKSKKHL	TDNEFK
	160	170	180	190	200
VR1	DPETGKTCLLKAM	LNLHNGONDTI	LLLDVARKTI	DSLKOFVNAS	YTDSYY
hVR1	DPETGKTCLLKAM	LNLHDGQNTTI	PLLLEIAROTI	SLKELVNAS	YTDSYY
	210	220	230	240	250
VR1	KGQTALHIATERR	NMTLVTLLVENG	ADVQAAANGI	FFKKTKGRP	GFYFGE
hVR1	KGQTALHIAIERR	NMALVTLLVENC	ADVQAAAHGI	FFKKTKGRP	GFYFGE
	260	270	280	290	300
VR1	LPLSLAACTNOLA	IVKFLLQNSWQI	PADISARDSVO	NTVLHALVE	VADNTV
hVR1	LPLSLAACTNOLG	IVKFLLONSWOI	ADISARDŠVO	NTVLHALVE	VADNTV
	310	320	330	340	350
VR1	DNTKFVTSMYNEI	LILGAKLHPTLE	LEEITNRÄGI	TPLALAASS	GKIGVL
hVR1	DNTKFVTSMYNEI	LILGAKLHPTLE	LEELTNKKGN	TPLALAAGT	GKIGVL
	360	370	380	390	400
VR1	AYILQREIHEPEC	RHLSRKFTEWAY	GPVHSSLYDI	SCIDTCEKN	SVLEVI
hVR1	AYILQREIQEPEC	RHLSRKFTEWAY	GPVHSSLÝĎÌ	SCIDTCEKN	SVLEVI
	410	420	430	440	450
VR1	AYSSSETPNRHDM	LLVEPLNRLLQL	KWDRFVKRIE	YFNFFVYCL	YMIIFT
	2314 C 24 5 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	and the second s	THE PARTY OF THE P	ART STREET SAN LINE AT THE PLAN.	The second second
VR1 hVR1	AYSSSETPNRHDM 460	LLVEPLNRLLQD 470	KWDREVKRIE 480	YENELVYCL 490	YMIIFT 500
	AYSSSETPNRHDM	LLVEPLNRLLQD 470	KWDREVKRIE 480	YENELVYCL 490	YMIIFT 500
hVR1 VR1	AYSSSETPNRHDM 460 AAAYYRPVEGLPP	LLVEPLNRLLOD 470 YKLKWIVGDYFF	KWDREVKRIE 480 VTGEILSVSG	YENELVYCL 490 GVYFFFRGI	YMLIFT 500 QYFLQR
hVR1	AYSSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510	LLVEPLNRLLOL 470 YKLKWIVGDYFF FKMEK . IGDYFF 520	KWDRFVKRIE 480 VTGEILSVS VTGEILSVLO 530	YFNFLVYCL 490 GVYFFERGI GVYFFFRGI 540	MILET 500 OYFLOR OYFLOR 550
hVR1 VR1	AYSSSETPNRHDM 460 AAAYYRPVEGTPP MAAYYRPVDGLPP	LLVEPLNRLLOL 470 YKLKWIVGDYFF FKMEK . IGDYFF 520	KWDRFVKRIE 480 VTGEILSVS VTGEILSVLO 530	YFNFLVYCL 490 GVYFFERGI GVYFFFRGI 540	MILET 500 OYFLOR OYFLOR 550
hVR1 VR1 hVR1 VR1	AYSSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS	LLVEPLNRLLOD 470 YKLKWTVGDYFF FKMEK . IGDYFF 520 EILFEYOSLFMI	OKWDREVKRIE 480 VTGEILSVSC VTGEILSVLC 530 VSVVLYESOE	YFNFLVYCL 490 GVYFFFRGI 540 KEYVASMVF	YMIIFT 500 OYFLOR OYFLOR 550 SLAMGW
hVR1 VR1 hVR1	AYSSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510	LLVEPLNRLLOD 470 YKLKWTVGDYFF FKMEK . IGDYFF 520 EILFEYOSLFMI	OKWDREVKRIE 480 VTGEILSVSC VTGEILSVLC 530 VSVVLYESOE	YFNFLVYCL 490 GVYFFFRGI 540 KEYVASMVF	YMIIFT 500 OYFLOR OYFLOR 550 SLAMGW
hVR1 VR1 hVR1 VR1	AYSSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS	LLVEPLNRLLOL 470 YKLKWIVGDYFF FKMEK . IGDYFF 520 EILFFYOSLFMI EMLFFLÖSLFMI 570	OKWDREVKRIE 480 VTGEILSVSC VTGEILSVLC 530 VSVVLYFSOF ATVVLYFSHI 580	YENFLVYCL 490 GVYFFFRGI GVYFFFRGI 540 KEYVASMVF KEYVASMVF	YMITET 500 OYFLOR 550 SLAMGW SLALGW 600
hVR1 VR1 hVR1 VR1 hVR1	AYSSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMRTLFVDSYS 560 TNMLYYTRGFOOM	LLVEPLNRLLQC 470 YKIKWIVGDYFF FKMEK IGDYFF 520 EILFFVOSLFMI EMLFFLÖSLFMI 570 GLYAVMIEKMII	OKWDREVKRIE 480 VTGEILSVSC VTGEILSVLC 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCRFMEVY	YENFLVYCL 490 GVYFEFRGI 540 KEYVASMVF KEYVASMVF 590 LVFLEGFST	YMILET 500 OYFLOR 550 SYFLOR 550 SLAMGW 600 AVVILI
hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMRTEFVDSYS 560	LLVEPLNRLLOD 470 YKLKWTVGDYFF FKMEK . IGDYFF 520 EILFFVOSLEMI 570 GIYAVMIEKMII GIYAVMIEKMII	AWDREVERIE 480 VTGEILSVLO 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVY	YFNFLVYCL 490 GVYFFFRGI GVYFFFRGI 540 RKEYVASMVF EYVASMVF 590 LVFLFGFST	YMILET 500 OYFLOR OYFLOR 550 SLAMGW 600 AVVILI AVVILI
hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMRTEFVDSYS 560 TNMLYYTRGFOOM TNMLYYTRGFOOM 610	LLVEPLNRILQI 470 YKLKWIVGDYFF FKMEK . IGDYFF 520 EILFFYOSLFMI 570 GIYAVMIEKMII 620	CWDREVERIE 480 VTGEILSVEC 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVY RDLCRFMFVY 630	490 GVYFFFRGI GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST	YMILET 500 OYFIOR 550 SIAMGW SLALGW 600 AVVILI 650
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNREDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTEFVDSYS 560 TNMLYYTRGFQOM TNMLYYTRGFQOM 610 EDGKNNSLPMEST	LLVEPLNRILOR 470 YKLKWTVGDYFF FKMEK.IGDYFF 520 EILFFYOSLFMI 570 GIYAVMIEKMII GIYAVMIEKMII 620 PHKCRGSACK.F	AWDREVERIE 480 VTGEILSVLO 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCREMEVY 630 GNSYNSLYST	YENFLVYCL 490 GVYFFERGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI	YMILET 500 OYFLOR OYFLOR SIANGW 600 AVVILL 650 ENGDLE
hVR1 VR1 hVR1 VR1 hVR1 VR1	AYSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS 560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST	LLVEPLNRILOR 470 YKLKWIVGDYFF 520 EILFEVOSLEMI EMLFELÖSLEMI 670 GIYAVMIEKMII 61YAVMIEKMII 620 PHKCRGSACK.F	AWDREVERTE 480 VTGEILSVLO 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMEVY 630 GNSYNSLYST	490 GVYFFERGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI	YMILET 500 OYFIOR 550 SLAMGW SLALGW 600 AVVILI 650 SMGDLE
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNREDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTEFVDSYS 560 TNMLYYTRGFQOM TNMLYYTRGFQOM 610 EDGKNNSLPMEST	LLVEPLNRILOR 470 YKLKWTVGDYFF 520 EILFEVOSLEMI 570 GIYAVMIEKMII 620 PHKCRGSACK.F	AWDREVERIE 480 AVTGEILSVLO 530 AVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVY 630 CGNSYNSLYST DSSYNSLYST 680	490 GVYFFERGI GVYFFFRGI STAND KEYVASMVF LKEYVASMVF 190 LVFLFGFST 640 CLELFKFTI 690	YMILET 500 OVELOR OVELOR STANGW SLANGW 600 AVVILL 650 SMGDLE MGDLE 700
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS 560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFRAVFII	LLVEPLNRILOR 470 YKLKWTVGDYFF 520 EILFEVOSLEMI EMLFELÖSLEMI GIYAVMIEKMII GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLCAYVILTYII	AWDREVERTE 480 VTGEILSVLG 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCREMEVY 630 GNSYNSLYST 680 LLIMMITALMG	490 GVYFFFRG1 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFT1 690 ETVNKTAOE	YMILET 500 OYFLOR OYFLOR SLANGW SLALGW 600 AVVTLI 650 EMGDLE 700 EKNIWK
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNREDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS 560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFKAVFIL	LLVEPLNRILOL 470 YKLKWIVGDYFF FKMEK.IGDYFF 520 EILFEVOSLFMI 570 GIYAVMIEKMII GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLAYVILTYII	AWDREVERTE 480 VTGEILSVLG 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMEVY 630 GNSYNSLYST 680 LLIMMLIALMG	YFNFLVYCL 490 GVYFFFRGI 540 KEYVASMVF KEYVASMVF 590 LVFLEGFST 640 CLELFKFTI 690 ETVNKTAOE	YMILET 500 OYFIOR 550 SLAMGW SLALGW 600 AVVILI 650 SMGDLE 700 SKNIWK
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS 7560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFKAVFII 710	LLVEPLNRILQL 470 YKLKWIVGDYFF FKMEK.IGDYFF 520 EILFFVOSLFMI 570 GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLAYVILTYII 720	OKWDREVKRIE 480 VTGEILSVSG 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCRFMFVV RDLCRFMFVV 630 GNSYNSLYST 080 LLNMLIAIMG 730	YENFLVYCL 490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740	YMILET 500 OYFIOR 550 SLAMGW SLALGW 600 AVVILI 650 SMGDLE MGDLE MGDLE FMGDLE FM
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1	AYSSETPNREDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS FOO TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFKAVFII FTENYDFKAVFII 710 LORALTILDTEKS	LLVEPLNRILOR 470 YKLKWTVGDYFF 520 EILFEVOSLEMI EMLFELOSLEMI 570 GIYAVMIEKMII 620 PHKCRGSACK.E SHRWRGPACRPE 670 LLTAYVILTYII LTAYVILTYII 720 ELKCMRKAFRSG	AWDREVERTE 480 VTGEILSVLG 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCREMEVY 630 GNSYNSLYST 680 LLNMLTALMG 730 KLLOVGETPE	490 GVYFFERGI 540 RKEYVASMVF KETVASMVF 100 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GKDDYRWGF	YMILET 500 OYFLOR OYFLOR SYFLOR SLANGW SLALGW 600 AVVILI 650 EMGDLE 700 EMGDLE 700 EKNIWE FROM
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNREDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLEVDSYS RPSMKTLEVDSYS 560 TNMLYYTRGEOOM TNMLYYTRGEOOM TNMLYYTRGEOOM EDGKNNSLEMEST EDGKNDSLPSEST 660 FTENYDEKAVELL 710 LORALTILDTEKS	LLVEPLNRILOL 470 YKLKWTVGDYFF FKMEK.IGDYFF 520 EILFEVOSLEMI 570 GIYAVMIEKMII GIYAVMIEKMII GIYAVMIEKMII GEO PHKCRGSACK.E SHRWRGPACRPE 670 LLLAYVILTYII 720 FLKCMRKAFRSG	AWDREVERTE 480 VTGEILSVS 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCRFMEVY 630 GNSYNSLYST 680 LLIMMLTAIMG 730 KLLOVGETPE	490 GVYFFERGI 540 KEYVASMVF X590 LVFLEGFST 640 CLELFKFTI 690 ETVNKTAOE 740 GKDDYRWCFI	YMILET 500 OYFIOR 550 SLAMGW SLALGW 600 AVVILI 650 ENGDLE 700 SKNIWK 750 KVDEVN
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMRTEFVDSYS 560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFRAVFIL 710 LORAITILDTEKS 760	LLVEPLNRILQL 470 YKLKWIVGDYFF FKMEK.IGDYFF 520 EILFFVOSLFMI 570 GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLAYVILTYII LLAYVILTYII LLAYVILTYII FLKCMRKAFRSG FLKCMRKAFRSG	AWDREVERTE 480 VTGEILSVSG 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVV 630 GNSYNSLYST DSSYNSLYST 1LINMLIALMG KLLOVGFTPL 780	YENFLVYCL 490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GRDDYRWCFI 790	YMILET 500 OYFIOR 550 SLAMGW SLALGW 600 AVVILI 650 EMGDLE FMGDLE
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1	AYSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS 7560 TNMLYYTRGFOOM TNMLYYTRGFOOM 610 EDGKNNSLPMEST 660 FTENYDFRAVFII FTENYDFRAVFII TORAITILDTEKS 760 WTTWNTNVGLINE	LLVEPLNRILQL 470 YKLKWTVGDYFF FKMEK.IGDYFF 520 EILFFYOSLFMI 570 GLYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPF 670 LLAYVILTYII T20 FLKCMRKAFRSG 770 DEGNCEGVKRTI	AWDREVERTE 480 VTGEILSVS 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVY RDLCRFMFVY 630 GNSYNSLYST 680 LLNMLTAIMG 730 KLLOVGFTPI 780 SFSLRSCRVS	490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GKDDYRWCFI 790 GRNWKNFAL	YMILET 500 OYFIOR 550 SIAMGW SLALGW 600 AVVILI 650 SMGDLE 700 SKNIWK 750 KNIWK 750 KVDEVN 800 PPLERD
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS 560 TNMLYYTRGFOOM TNMLYYTRGFOOM TNMLYYTRGFOOM EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFKAVFII 710 LORAITILDTEKS 760 WITWNTNVGLINE	LLVEPLNRILOL 470 YKLKWTVGDYFF FKMEK IGDYFF 520 EILFEVOSLEMI 570 GIYAVMIEKMII GIYAVMIEKMII GIYAVMIEKMII GIYAVMIEKMII 620 PHKCRGSACK F SHRWRGPACRPE 670 LLLAYVILTYII 720 FLKCMRKAFRSG 770 DPGNCEGVKRTI	AWDREVERTE 480 VTGEILSVS 530 VSVVLYFSOE ATVVLYFSHI 580 RDLCRFMEVY 630 GNSYNSLYST 680 LLIMMLTAIMG 730 KLLOVGFTPI 780 SFSLRSSRVS	490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GKDDYRWCFI 790 GRNWKNFAL	YMILET 500 OYFIOR 550 SIAMGW SLALGW 600 AVVILI 650 SMGDLE 700 SKNIWK 750 KNIWK 750 KVDEVN 800 PPLERD
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1	AYSSETPNRHDM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMRTEFVDSYS 560 TNMLYYTRGFOOM 610 EDGKNNSLPMEST 660 FTENYDFRAVFIL 710 LORAITILDTEKS 760 WITWNTNVGLINE WITWNTNVGLINE 810	LLVEPLNRILQE 470 YKLKWIVGDYFF FKMEK.IGDYFF 520 EILFEVOSLFMI 570 GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLAYVILTYII LLAYVILTYII LLAYVILTYII CHARKAFRSG FLKCMRKAFRSG FLKCMRKAFRSG FLKCMRKAFRSG PGNCEGVKRTI DPGNCEGVKRTI	AWDREVERTE 480 VTGEILSVS 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVY 630 GNSYNSLYST OSSYNSLYST ALLNMLIALMO KLLOVGFTPL 780 SFSLRSGRVS 830	YENFLVYCL 490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GRDDYRWCFI 790 GRNWKNFALV	YMILET 500 OYFIOR 550 SIAMGW SLALGW 600 AVVILI 650 SMGDLE 700 SKNIWK 750 KNIWK 750 KVDEVN 800 PPLERD
hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1 hVR1 VR1	AYSSETPNRHOM 460 AAAYYRPVEGLPP MAAYYRPVDGLPP 510 RPSLKSLFVDSYS RPSMKTLFVDSYS 560 TNMLYYTRGFOOM TNMLYYTRGFOOM TNMLYYTRGFOOM EDGKNNSLPMEST EDGKNDSLPSEST 660 FTENYDFKAVFII 710 LORAITILDTEKS 760 WITWNTNVGLINE	LLVEPLNRILQI 470 YKLKWIVGDYFF FKMEK.IGDYFF 520 EILFFYOSLFMI 570 GIYAVMIEKMII 620 PHKCRGSACK.F SHRWRGPACRPE 670 LLAYVILTYII 720 FLKCMRKAFRSG FLKCMRKAFRSG FLKCMRKAFRSG 100 PGNCEGVKRTI 820 VOLKHYTGSLKE	CWDREVERIE 480 VTGEILSVS 530 VSVVLYFSOF ATVVLYFSHI 580 RDLCRFMFVV RDLCRFMFVV 630 GNSYNSLYST 080 LLNMLIAIMO 730 KLLOVGFTPF 780 SFSLRSGRVS SFSLRSGRVS 830	490 GVYFFFRGI 540 KEYVASMVF 590 LVFLFGFST 640 CLELFKFTI 690 ETVNKTAGE 740 GKDDYRWCFI 790 GRNWKNFALV	YMILET 500 OYFIOR 550 SIAMGW SLALGW 600 AVVILI 650 SMGDLE 700 SKNIWK 750 KNIWK 750 KVDEVN 800 PPLERD

FIG. 17

hVR3 SEQUENCE INCLUDING 5' UTR (nt -686 TO nt 0) CODING REGION (nt1 TO nt 2889), 3'UTR (nt 2890 TO nt 3418)--(SEQ ID NO:4)--

-684	ttacgcgttaagaaatacccaagcttatgcatcaagcttggtaccgagctcggatccact	-625
-624	agtaccgccggccagtgtgctggaattcaaggtgaggaggaggagcatggatcctgggagc	-565
-564	gagtgtgtgcaggccagggagggctttccagaggagcccagttgagctggaacaccagtg	-505
-504	gggaggagttgaccagcaaaggtgcagggagggatcagcactttgcactggggagcagag	-445
444	tttgtgcactggggaagtcaactcaagtattggagcctcagtttcctgttctgtaaaatg	-385
384	ggttcatcatgacagtgtttgatgaggaaaaggactgccggcctacacagcaagtccaca	-325
324	tggattttctgagcccctcctgtgcctgaagcccacggttaatggttctgccttagcagg	-265
264	tgettaccacgtgccaggcactgcactgcactggccactggactgcatgttctgtccatg	-205
204	aggettggatatececatettacagateaggaagetgaggetatgaaatgtegaettget	-145
144	caatgteatggaatgaetaagtgtggageetggatttgaaettggetetetggggeteea	-85
-84	aagctggctttcttggtcagcagtagggtctgggatccaagtatggggtcccagcttgac	-25
-24	cctgaagtccaccctctttcagctaATGCCCAGGGTAGTTGGACCTGGGGCCAATTTGTG	35
36	TTTCCAGGTTCGTGAAAGAGGCTCCTGTTGCAGTTCCCGCCTGAGGCTGGCGGCCAACCA	95
96	CATCTGGGAGTGGCCTCCCTGTGCCCCTGTCATTACAACGGTGGCTTTGAAGCAGCTGGC	155
156	AGCÁCTGCTGCTTGTCCACGTGGGAGGGGGCTTCCTGGAGCCCCCGCCCCTGGCCGGGTT	215
216	CTGCCTGACTCCCCTTTCATTCCCTTGCAGGCTGAGCAGTGCAGACGGGCCTGGGGCAGG	275
276	CATGCCGGATTCCAGCGAAGGCCCCCGCGCGGGGGCCCCGGGGGGGG	335
336	CC N MC N C N C C C C N C C C C C C C C	305

FIG. 18

NUCLEOTIDE AND AMINO ACID SEQUENCE OF hVR3 --(SEQ ID N0:4)-INCLUDING THE 5'UTR (nt -684 TO nt 0), CODING REGION (nt1
TO 2889) AND 3'UTR (nt 2890 TO nt 3418)--(SEQ ID N0:5)--

-684	ttacgcgttaagaaatacccaagcttatgcatcaagcttggtaccgagctcggatccact	-625
-624	agtaccgccggccagtgtgctggaattcaaggtgaggaggaggatggat	-565
-564	gagtgtgtgcaggccagggagggctttccagaggagcccagttgagctggaacaccagtg	-505
-504	gggaggagttgaccagcaaaggtgcagggagggatcagcactttgcactggggagcagag	-445
-444	tttgtgcactggggaagtcaactcaagtattggagcctcagtttcctgttctgtaaaatg	-385
-384	ggttcatcatgacagtgtttgatgaggaaaaggactgccggcctacacagcaagtccaca	-325
-324	tggattttctgagcccctcctgtgcctgaagcccacggttaatggttctgccttagcagg	-265
-264	tgcttaccacgtgccaggcactgcactgcactggccactggactgcatgttctgtccatg	
-204	aggettggatatececatettacagateaggaagetgaggetatgaaatgtegaettget	
144	caatgtcatggaatgactaagtgtggagcctggatttgaacttggctctctggggctcca	-85
-84	aagetggetttettggteageagtagggtetgggateeaagtatggggteeeagettgae	-25
-24 1	cctgaagtccaccctctttcagctaATGCCCAGGGTAGTTGGACCTGGGGCCAATTTGTG M P R V V G P G A N L C	35 12
36	TTTCCAGGTTCGTGAAAGAGGCTCCTGTTGCAGTTCCCGCCTGAGGCTGGCGGCCAACCA	95
13	FQVRERGSCCSSRLRLAANH	32
96	CATCTGGGAGTGGCCTCCTGTGCCCCTGTCATTACAACGGTGGCTTTGAAGCAGCTGGC	155
33	IWEWPPCAPVITTVALKQLA	52
156	AGCACTGCTGCTGCACGTGGGAGGGGGGCTTCCTGGAGCCCCCGCCCCTGGCCGGGTT	215
53	ALLLVHVGGGFLEPPLAGF	72
216	CTGCCTGACTCCCCTTCATTCCCTTGCAGGCTGAGCAGTGCAGACGGCCTGGGGCAGG	275
73	C L T P L S F P C R L S S A D G P G A G	92
276	CATGGCGGATTCCAGCGAAGGCCCCGCGCGGGGGCCCGGGGAGGTGGCTGAGCTCCCCGG	225
93	M A D S S E G P R A G P G E V A E L P G	335 112
226		112
336	GGATGAGAGTGGCACCCCAGGTGGGGAGGCTTTTCCTCTCTCT	395
113	DESGTPGGEAFPLSSLANLF	132
396	TGAGGGGGAGGATGGCTCCCTTTCGCCCTCACCGGCTGATGCCAGTCGCCCTGCTGGCCC	455
133	E G E D G S L S P S P A D A S R P A G P	455 152
		132
456	AGGCGATGGCCGACCAAATCTGCGCATGAAGTTCCAGGGCGCCCTTCCGCAAGGGGGTGCC	515
153	GDGRPNLRMKFQGAFRKGVP	172
516	CAACCCCATCGATCTGCTGGAGTCCACCCTATATGAGTCCTCGGTGGTGCCTGGGCCCAA	575

FIG. 19

AMINO ACID SEQUENCE OF hVR3 -- (SEQ ID NO:5) --

MPRVVGPGAN LCFQVRERGS CCSSRLRLAA NHIWEWPPCA PVITTVALKQ LAALLLVHVG GGFLEPPPLA GFCLTPLSFP CRLSSADGPG AGMADSSEGP RAGPGEVAEL PGDESGTPGG EAFPLSSLAN LFEGEDGSLS PSPADASRPA 101 GPGDGRPNLR MKFQGAFRKG VPNPIDLLES TLYESSVVPG PKKAPMDSLF DYGTYRHHSS DNKRWRKKII EKQPQSPKAP APQPPPILKV FNRPILFDIV 201 SRGSTADLDG LLPFLLTHKK RLTDEEFREP STGKTCLPKA LLNLSNGRND 251 TIPVLLDIAE RTGNMREFIN SPFRDIYYRG QTALHIAIER RCKHYVELLV 301 AQGADVHAQA RCRFFQPKDE GGYFYFGELP LSLAACTNQP HIVNYLTENP 351 HKKADMRROD SRGNTVLHAL VALADNTREN TKFVTKMYDL LLLKCARLFP 401 DSNLEAVINN DGLSPIMMAA KTGKIGIFQH IIRREVTDED TRHISRKSKD 451 WAYGPVYSSL YDLSSLDTCG EEASVLEILV YNSKIENRHE MLAVEPINEL 501 LRDKWRKFGA VSEYINVVSYLCAMVIETLTEAYYOPLEGTP PYPYRTTVDY 551 LRLAGEVITETETGVIDEFETN IKDLEMKKCP GVNSLFIDGS: FOLLYFIYSV 601 ININSAALYL AGIEAYLAMM VFALVLGWMN ALYETRGLKLETGTYSIMEQK 651 ILFKDI FRETERIAMMUEMIGYSASAMSLINP CANMKVCNED QTNCTVPTYP SCRDSETFST FLLDLFKLTI CMCDLEMLSS TKYPWFIIL LVTYIILTSV 751 LILINMIHALMEGETVGQVSKE SKHIWKLQWA TTILDIERSF PVFLRKAFRS 801 GEMVTVGKSS DGTPDRRWCF RVDEVNWSHW NQNLGIINED PGKNETYQYY 851 GFSHTVGRLR RDRWSSVVPR VVELNKNSNP DEVVVPLDSM GNPRCDGHQQ 901 GYPRKWRTDD APL 951

Key

Transmembrane domains

Ankyrin binding domains

FIG. 21

A MULTIPLE COMPARISON OF THE AMINO ACID SEQUENCES OF THE RAT VR1 AND THE HUMAN VANILLOID RECEPTORS, hVR1, hVRL-1 AND hRV3

		10	20	30	40	50			0)
VR1	~~~~~~	~~~~~~	20	~~~~~~		~~~	(SEQ	ID	NO:3)
hVR1	~~~~~	~~~~~~	~~~~~~	~~~~~~		~~~	(SEQ	ID	NO:2)
hVRL-1	~~~~~~	~~~~~~~	~~~~~~				(SEQ	ID	NO:6)
hVR3	MPRVVGPG.	ANLCFQVRER	GSCCSSRLRL		PCAPVITTVA	TKÖ	(SEQ	ID	NO:5)
		60	70	80	90	100			
VR1	~~~~~		~~~~~~						
hVR1		~~~~~~	~~~~~~	~~~~~~		~~~			
hVRL-1 hVR3		vcccet.eddd	LAGFCLTPLS	EDCDT.GGAD	PCACMADSS	ECD			
HVKS	THAT DE VII								
VR1		110	120	130	140	150			
hVR1			~~~~~~						
hVRL-1			~~~~~~						
hVR3	RAGPGEVA	ELPGDESGTP	GGEAFPLSSI	ANLFEGEDG	SLSPSPADAS	RPA			
		160	170	180	190	200			
VR1			FTTRSKTŔĹ						
hVR1			STAKSRTRÎÎ						
hVRL-1	~~~~~~~ ~Ef ~20 ~ 1275~~	~~~~~~	MTSPSSSPVE	RLETLDGGQI	EDGSEADRGK	LDF			
hVR3	GÄGÖGÉFA		KGVPNP						
		210	220	230	240	250	-		
VR1		Carried and the second and the second	PGDGPASVŘÍ PGDGPTGARI	and the second second	which is the property of the contract of the c	3-2-3-3			
hVR1 hVRL-1			KFAPQIRVNI						
hVR3			RWRKKIIEK						
		260	270	280	290	300			
VR1	TEDAVAOS		FLORSKKRLI						
hVR1			FLOKSKKHLI						
hVRL-1	LENAVSRG	VPEDLAGLPE	YLSKTSKYL	DSEYTEGST	GKTCLMKAVI	NEK			
hVR3	LEDIVSRG	STADLDGLLL	FLLTHKKRLT	DEEFREPST	GKTCLPKALI	NUS			
		310 .	320	330	340	350			
VR1			SLKOFVNAS						
hVR1	DCONTILE	LLLE LAROTI	SLKELVNASI	TDSYYKCOL	ALHIAIERRN	MALI			
hVRL-1 hVR3	DGVNACIL	PLIQIDEDSC	npoplvnaoc NMREFINSPI	TDDXXRGHS	ALHIAIEKKS	PLOC			
IIVKS	MGRADITE								
		360	370 FFKKTKGKP	380	390	400			
VR1 hVR1			FFKKTKGRP						
hVRL-1	VKLLVENG	ANVHARACGE	CFOKGOG TO	EYEGELPLS	LÄACTKOWDV	V SY			
hVR3	VELLVAQG	ÄDVHAQARĞI	VFFQPKDEGGY	FYFGELPLS	LAACTNOPHI	YNY			
		410	420	430	440	450			
VR1	HEINON SINOS	ADTSARDSVO	NTVLHAEVEY	VADINUA VIDNIEK	EVTSMYNE II	TE			
hVR1	TOWSMOT	ADISARDSVO	ntvlhälvet	ADNIVADNIK	fytsmyne ii	HG			
hVRL-1	TEENPHOR	ASLOATOSO	NTVĽHALVMI	SDNSAENIA	LVTSMYDGLI	QAG			
hVR3	LTENPHKK	ADMRRODSRO	NTVLHALVA	LADNTRENTK	EALKWADITI	TKC			
_		460	470	480	490	500			
VR1	AKEHETEK	LEE I TNRKGI	TPLALASS	KIGVLAYIL	ORE THE PECI	CHIES:			
hVR1 hVRL-1	AVTH PATE	PER LEGIT OF	itplaľaagt Tplkľaake	SELECTED LET	VARIATION CO				
hVR3	ARDEPOSN	LEAVINDG	SPIMMAKT	KIGIFOHII	RREVIDEDI				
					hapter 7				